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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

C08L 23/10, 23/08, 23/04, C08J 5/18, B32B 27/32

A1

(11) International Publication Number:

WO 98/37139

(43) International Publication Date:

27 August 1998 (27.08.98)

(21) International Application Number:

PCT/US98/03597

(22) International Filing Date:

24 February 1998 (24.02.98)

(30) Priority Data:

25 February 1997 (25.02.97) US 08/806,182 28 April 1997 (28.04.97) US 08/847,730 08/905,211 US

1 August 1997 (01.08.97)

(63) Related by Continuation (CON) or Continuation-in-Part (CIP) to Earlier Applications

08/806,812 (CIP) US Filed on Not furnished 08/847,730 (CIP) US Not furnished Filed on 08/905,211 (CIP) US Not furnished Filed on

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(81) Designated States: AU, BR, CA, CN, HU, IL, JP, KR, NO, NZ, PL, PT, SG, TR, US, European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).

Published

With international search report.

(54) Title: HEAT SEALABLE FILMS

(57) Abstract

This invention relates to a film comprising a blend comprising: (i) a homopolymer of ethylene having an Mw/Mn of 3 or less or a copolymer of ethylene and up to 50 weight % of a C3 to C20 olefin having a CDBI of 50 % or more, (ii) a homopolymer of propylene or copolymer of propylene and up to 50 weight % of a comonomer, and (iii) a polymer produced in a high pressure process using a free radical initiator. In a preferred embodiment the blend comprises mPE, LDPE and isotactic PP.

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HEAT SEALABLE FILMS

Field of the Invention

5 This invention relates to films comprising a blend comprising metallocene polyethylene (mPE), polypropylene and free radically high pressure produced polymers.

Background of the Invention

- Polyethylene has traditionally been used as a layer in polyolefin films and 10 packaging because of its desirable properties such as moisture impermeability good sealing behavior, good optical properties and good organoleptics. Typically polyethylene has been coextruded, laminated or otherwise bonded to other polyolefins which have better strength than polyethylene, yet do not seal as well as polyethylene. For example, in a typical multilayer film, a polypropylene layer. 15 especially a mono or biaxially oriented polypropylene (OPP) layer, provides a high clarity, high melting, high barrier properties, combined with high stiffness, while a polyethylene layer will provide extra body to the film and will allow a low : sealing temperature, meaning higher packaging speeds. However, polypropylene : (PP) and polyethylene (PE) have very limited compatibility and direct sealing of 20 polyethylene onto polypropylene film is not commonly done. When a layer of PE is combined with a layer of PP, extra primer may be needed. For example, extra
- onto polypropylene films. In addition tie layers may also be necessary.

 Coextrudable tie layers such as ethylene vinyl acetate copolymers, typically having more that 8 weight % vinyl acetate, have been extruded between PP and PE to enhance adhesion between the PE and the PP. Another solution to the compatibility problem has been to blend polypropylene into the polyethylene. This however has the disadvantage or creating layers that have greater haze and are thus undesirable in the industry.

primer is used when polyethylene, such as low density polyethylene, is coated

Therefore there is a need in the art to provide a means to provide a polyethylene polypropylene blend for film layers that does not have haze, yet retains good mechanical properties such impact strength.

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WO 94/26816 discloses blends of metallocene polyethylene and high molecular weight high density polyethylene for use in films.

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Summary of the Invention

This invention relates to a film comprising a blend of:

(i) a homopolymer of ethylene having an Mw/Mn of 3 or less or a copolymer of 5 ethylene and up to 50 weight % of a C₃ to C₂₀ olefin, wherein the copolymer has a CDBI of 50 % or more, preferably 60% or more;

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- (ii) a homopolymer of propylene or a copolymer of propylene and up to 50 weight% of a comonomer, preferably copolymerized with ethylene and/or a C₄ to C₂₀ Constitution of the contract o
- 10 (iii) a polymer produced in a high pressure process using a free radical initiator (High Pressure Polymer). The appeals strained as the contract of straining of

This invention also relates to films as described above where one or more layers are oriented in one or more directions to the same or different extents.

Detailed Description of the Invention and the second secon

In a preferred embodiment, this invention relates to a film comprising a blend of:

- (i) a homopolymer of ethylene having an Mw/Mn of 3 or less, preferably between 1 and 2.5 or a copolymer of ethylene and up to 50 weight %, preferably 1 to 35 20 weight %, preferably 1-20 weight % of one or more C₃ to C₂₀ olefins, (based upon the weight of the copolymer) having an Mw/Mn of 6 or less, preferably 3 or less, even more preferably between 1 and 2.5, wherein the polymer or copolymer preferably has: $\frac{1}{2}$ $\frac{1}$
- more preferably between 0.88 g/cm² and 0.935 g/cm³, more preferably between 0.88 g/cm³ and 0.95 g/ cm³, more preferably between 0.915 g/cm³; and 0.935 g/ cm³; and
 - b) a CDBI of 50 % or more, preferably above 60%;
 - (ii) a homopolymer of propylene or a copolymer of propylene and up to 50 weight %, preferably 1 to 35 weight %, even more preferably 1 to 6 weight % of ethylene and/or a C4 to C20 olefin; and a grant of the least of the case of th
- 30 . Fig., (iii) and polymer produced in a high pressure process using a free radical initiator The transfer of the High Pressure Polymer). The design to the design of the second to the second second the second second

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Composition Distribution Breadth Index (CDBI) is a measure of the composition distribution of monomer within the polymer chains and is measured by the procedure described in PCT publication WO 93/03093, published February 18, 1993 including that fractions having a weight average molecular weight (Mw) below 15,000 are

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ignored when determining CDBI. For purposes of this invention a homopolymer is defined to have a CDBI of 100%.

The C₃ to C₂₀ and C₄ to C₂₀ olefin comonomers for the polyethylene or polypropylene copolymers described above may be any polymerizable olefin monomer and are preferably a linear, branched or cyclic olefin, even more preferably an α-olefin. Examples of suitable olefins include propylene, butene, isobutylene, pentene, isopentene, cyclopentene, hexene, isohexene, cyclohexene, heptene, isohexene, isohexene, sohexene, decene, isohexene, cyclononene, decene, isodecene, dodecene, isodecene, decene, isodecene, trienes, and styrenic monomers.

Preferred examples include styrene, α-methyl styrene, para-alkyl styrene (such as para-alkyl styrene), hours linear para-alkyl styrene (such as para-alkyl styrene).

methyl styrene), hexadiene, norbornene, vinyl norbornene, ethylidene norbornene, butadiene, isoprene, heptadiene, octadiene, and cyclopentadiene.

Preferred comonomers for the copolymer of ethylene are propylene, butene, hexene and/or octene.

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termonomers and tetramonomers which may be one or more of the C3 to C20 olefins described above, any C4 to C30 linear, cyclic or branched dienes or trienes and any styrenic monomers such as styrene, or methyl styrene, or para-methyl styrene.

Preferred examples include butadiene, pentadiene, cyclopentadiene, hexadiene, cyclohexadiene, heptadiene, octadiene, nonadiene, norbornene, vinyl norbornene.

The polyethylene copolymers described above preferably have a composition distribution breadth index (CDBI) of 50 % of more, preferably above 60%, even more preferably above 70%. In one embodiment the CDBI is above 80%, even more preferably above 95%. In another particularly preferred embodiment, the polyethylene copolymer has a CDBI between 60 and 85 %, even more preferably between 65 and 85 %.

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In another preferred embodiment the ethylene homopolymer of copolymer has a density of 0.86 to 0.925 g/cm³ and a CDBI of over 80%, preferably between 80 and 99%.

In another preferred embodiment the blend comprises a homopolymer of ethylene having an Mw/Mn of 3 or less, preferably between 2.5 and 1.

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In general, the polyethylene homopolymers and copolymers described above are metallocene polyethylenes (mPE's). The mPE homopolymers or copolymers are typically produced using mono- or bis-cyclopentadienyl transition metal catalysts 10, in combination with an activator such as alumoxane and/or a non-coordinating anion in solution, slurry, high pressure or gas phase. The catalyst and activator may be supported or unsupported and the cyclopentadienyl rings by may substituted or unsubstituted. Several commercial products produced with such catalyst/activator combinations are commercially available from Exxon Chemical 15 Company in Baytown Texas under the tradenames EXCEED™ and EXACT™. 31 For more information on the methods and catalysts/activators to produce such mPE homopolymers and copolymers see WO 94/26816; WO 94/03506; EPA 277,003; EPA 277,004; US 5,153,157; US 5,198,401; US 5,240,894; US 5,017,714; CA 1,268,753; US 5,324,800; EPA 129,368; US 5,264,405; EPA .520,732; WO 92:00333; US 5:096,867; US 5,507,475; EPA 426 637; EPA 573 403; EPA 520,732; EPA 495,375; EPA 500,944; EPA 570,982; WO91/09882; WO94/03506 and US-5,055,438 dimension for the Automotive of

> The polypropylene homopolymer or copolymer preferably has an MFR (melt flow rate) of 1 to 20 as measured according to ASTM D 1238 (230 °C, 2.16 kg). In another embodiment the polypropylene homopolymer or copolymer preferably has a CDBI of 50 % or more, preferably above 60%, even more preferably above 70 %. Polypropylenes having a CDBI above 60% are available from Exxon Chemical Company in Baytown; Texas under the tradename ACHIEVETM

gradules as the mean Office of them of the managers by the some in the

In another embodiment the polypropylene homopolymer or copolymer can be blended with any of the other propylene homopolymers or copolmyers described above. Likewise, The polyethylene homopolymers or copolymers described above for use, in the blend may be used alone, may be blended with any of the other polyethylene homopolymers or copolymers described above.

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In a preferred embodiment the polymer produced in a high pressure process using a free radical initiator (High Pressure Polymer) is a polymer comprising one or more of C2 to C20 olefins and polar monomers. Preferred C2 to C20 olefins include, but are not limited to, ethylene, propylene, butene, pentene, hexene. octene, 3-methyl-pentene-1, 4-methyl-pentene-1, cyclopentene, cyclohexene, hexadiene, norbornene, isobutene, norbornadiene, pentadiene and 3.5.5-trimethyl hexene -1. Preferred polar monomers include, but are not limited to, acetates (such as vinyl acetate), acrylics (such as acrylic acid, methacrylic acid), acrylates (such as methacrylate, butylacrylate, methylmethacrylate, hoperated with the hydroxyethylmethylacrylate): Polar modifiers can also be used in high pressure

free radical process such as alcohols (such as isopropanol) or aldehydes (such as acetaldehyde). Other modifiers known in the art can also be used. Standards employed a colors, have expensed a some to compress and the standards.

In a preferred embodiment the High Pressure Polymer is low density polyethylene (density 0.910 to less than 0.940 g/cm³, preferably 0.915 to less than 0.935 g/cm³, Even more preferably 0.920 to less than 0.935 g/cm³), a copolymer of ethylene and vinyl acetate, a copolymer of ethylene and methyl acrylate, a copolymer of acrylic acid. a copolymer of methylmethacrylate or any other polymers polymerizable by a highpressure free radical process. The LDPE preferably has up to 20 weight % of comonomer. The EVA and acrylate copolymers preferably has 20 weight % of the polar monomer or less, preferably less than 10 weight %, even more preferably less than 6 weight %. In a preferred embodiment the Melt Index of the LDPE is between 0.2 and 50 g/10 min, preferably between 0.5 and 10 g/10 min, even more preferably between 0.6 and 5 g/10 min, even more preferably between 0.6 and 2.5 g/10 min.

Many such High Pressure Polymers are commercially available. For example, LDPE made in a high pressure process is available from Exxon Chemical Company under the trade name ESCORENE TM. EVA made in a high pressure process is available from Exxon Chemical Company under the trade name ESCORENETM.

Polymethylmethacrylate made in a high pressure process is available from Exxon Chemical Company under the trade name ESCORENETM 1996 218 300000 to the life of a stream or increasing among the contract of the second is a second

In a preferred embodiment the polyethylene (component (i)) is present in the 35 blend at from 1 to 99 weight %, based upon the weight of the polymers in the blend, preferably the polyethylene is present at 10 to 90 weight %, even more

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- In a preferred embodiment the polypropylene (component (ii)) is present in the blend at from 1 to 99 weight %, based upon the weight of the polymers in the blend, preferably the polypropylene is present at 10 to 90 weight %, even more preferably at least 20 to 80 weight %, even more preferably at least 30 to 70 weight %, even more preferably at least 40 to 70 weight %.
- In a preferred embodiment the High Pressure Polymer (component (iii)) is present in the blend at from 1 to 50 weight %, based upon the weight of the polymers in the blend, preferably at 2 to 30 weight %, even more preferably at least 5 to 20 weight %.

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- The blends described above can also further include other polymers such as polybutene, high density polyethylene (density 0.945 to less than 0.98 g/cm³) linear low density polyethylene, medium density polyethylene (density 0.935 to less than 0.945 g/cm³), polyyinylchloride, isotactic polybutene, ABS resins, elastomers such as ethylene-propylene rubber (EPR), vulcanized EPR, EPDM, block copolymer elastomers such as SBS, pylons, polygorbanetas, PET resins.
 - block copolymer elastomers such as SBS, nylons, polycarbonates, PET resins, crosslinked polyethylene copolymers of ethylene and vinyl alcohol (EVOH), polymers of aromatic monomers such as polystyrene; poly-1 esters, graft copolymers generally, polyacrylonitrile homopolymer or copolymers, thermoplastic polyamides, polyacetal, polyvinylidine fluoride and other
 fluorinated elastomers, polyethylene glycols and polyisobutylene.
- The blends described above may be produced by mixing the three (or more) polymers together, by connecting reactors together in series to make reactor blends or by using more than one catalyst in the same reactor to produce multiple species of polymer. The polymers can be mixed together prior to being put into an extruder or may be mixed or compounded in an extruder.

and the many commenced many and a significance.

The blends described above are typically formed into monolayer or multilayer films. These films may be formed by any of the conventional techniques known in the art including extrusion, co-extrusion, extrusion coating, lamination, blowing and casting. The film may be obtained by the flat film or tubular process which

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may be followed by orientation in an uniaxial direction or in two mutually perpendicular directions in the plane of the film and the second a

In a preferred embodiment a film of the blend is used as a sealing layer. In another preferred embodiment a film of the blend is used as that functional layer. that is to say it is used to provide mechanical strength and/or stiffness. Films of the blends described herein have excellent stiffness and mechanical strength as compared to films of the individual components.

- This invention also relates to films as described above where one or more of the layers are oriented in the transverse and/or longitudinal directions to the same or different extents. This orientation may occur before or after the individual layers are brought together. For example the blend layer can be extrusion coated or laminated onto another layer or the layers can be coextruded together into a film then oriented.
- 15 Typically the films are oriented in the Machine Direction (MD) at a ratio of up to 15, preferably between 5 and 7, and in the Transversé Direction (TD) at a ratio of up to 15 preferably 7 to 9. However in another embodiment the film is oriented to the same extent in both the MD and TD directions. Orientation to the same extent in both directions will generally produce roughly equal mechanical properties.

of the fifth of the control of the c In another embodiment the blend layer is combined with one or more other layers. The other layer(s) may be any layer typically included in multilayer film structures. For example the other layer or layers may be hard and part of the control of the cont The same of the manager forecast and the contract of

Polyolefins so the mass energy energiated and in the most 25 1. Preferred polyolefins include homopolymers or copolymers of C₂ to C₄₀ olefins, preferably C₂₀ to C₂₀ olefins, preferably a copolymer of an α-olefin and another olefin or α-olefin (ethylene is defined to be an α-olefin for purposes of this invention). Preferably homopolyethylene, homopolypropylene, propylene copolymerized with 30 ethylene and or butene, ethylene copolymerized with one or more of propylene, butene or hexene, and optional dienes. Preferred examples include thermoplastic polymers such as ultra low density polyethylene, very low density polyethylene, linear low density polyethylene, low density polyethylene, medium density polyethylene, high density polyethylene, polypropylene, isotactic polypropylene, highly isotactic 35 propolypropylene, syndiotactic polypropylene, random copolymer of propylene and ethylene and/or butene and/or hexene, elastomers such as ethylene propylene rubber, ethylene propylene diene monomer rubber, neoprene, and blends of thermoplastic

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polymers and elastomers, such as for example, thermoplastic elastomers and rubber Definition toughened plastics. From the large terms of the second of the

Polar polymers

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Preferred polar polymers include homopolymers and copolymers of esters, amides, 5 actates, anhydrides, copolymers of a C2 to C20 olefin, such as ethylene and/or propylene and/or butene with one or more polar monomers such as acetates, anhydrides, esters, alcohol, and or acrylics. Preferred examples include polyesters, polyamides, ethylene vinyl acetate copolymers, and polyvinyl chloride. 10 to Early to the notice of a crossed gardese score offer the engage with a con-

Cationic polymers

Preferred cationic polymers include polymers or copolymers of geminally disubstituted olefins, alpha-heteroatom olefins and/or styrenic monomers. Preferred geminally disubstituted olefins include isobutylene, isopentene, isoheptene, isohexane, isooctene, isodecene, and isododecene. Preferred alpha-heteroatom olefins include vinyl ether and vinyl carbazole, preferred styrenic monomers include styrene, alkyl styrene, para-alkyl styrene, alpha-methyl styrene, chloro-styrene, and bromo-para-methyl styrene. Preferred examples of cationic polymers include butyl rubber, isobutylene copolymerized with para methyl styrene, polystyrene, and poly-α-methyl styrene.

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and of another than the state of the state of the state of the state of Further any of the above layers may be oriented before or after being combined with the blend layers. 在1960年中中 · 大型10

A particularly preferred embodiment includes an ABC structure film where the A layer comprises mPE or a blend comprising mPE and the B layer is a blend according to this invention and the C layer is a sealing layer for example a random copolymer of propylene and up to 20 weight % of ethylene, preferably 3 to 6

35 weight % ethylene, even more preferably 3.5 to 5.5 weight % ethylene, or a At the sterpolymer of propylene, ethylene and butene. "我们不知知我们还要是有的数据,一个人的数据证明,这种类似的。" 经费 大部门的现在分词 医自己性神经

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In a preferred embodiment up to 100 µm thick monolayer films of the blend described above are characterized by a haze, as measured by ASTM 1003 condition A of 16% or less, more preferably 14% or less, more preferably 12 % or less, more preferably 10% or less, even more preferably 5% or less.

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The films described herein may vary in thickness depending on the intended application, however films of a thickness from 1 to 350 µm are usually suitable. Films intended for packaging are usually from 10 to 120 µm thick. The thickness of the sealing layer is typically 0.2 to 50 µm. There may be a sealing layer on both the inner and outer surfaces of the film or the sealing layer may be present on only the inner or the outer surface.

Additives such as antiblock, antioxidants, pigments, fillers, processing aids, UV mass translations, neutralizers, lubricants, surfactants and/or nucleating agents may also be present in one or more than one layer in the films. Preferred additives include silicon dioxide, titanium dioxide, polydimethylsiloxane, talc, dyes, wax, calcium stearate, carbon black, low molecular weightfresins, tackifiers, and glass beads.

In another embodiment the layers may be modified by corona treatment, electron beam irradiation, gamma irradiation, or microwave. In particular the corona treatment will produce a significant difference in the coefficient of friction of the two surface layers as described in US Patent Application Number USSN 08/905,211, which is incorporated by reference herein.

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In a particularly preferred embodiment film of the blends described herein are cast, blown or co-extruded and the polyethylene is present at 50 to 80 weight %, the polypropylene is present at 10 to 45 weight% and polymer produced in high pressure process using a free radical initiator is LDPE and is present at 2 to 10 weight%, based upon the weight of the blend.

In another embodiment films of the blends described herein are laminated to a substrate. Preferred substrates include polypropylene, polyamide, polyester, polyethylene, or metallized substrates.

The films described herein may also comprise from 5 to 60 weight %; based upon the weight of the polymer and the resin, of a hydrocarbon resin. The resin preferably has a softening point above 100 °C, even more preferably from 130 to 180 °C, even more

124	preferably between 140 and 180 °C. Preferred hydrocarbon resins include those
	described in EPA 288 227 and EPA 247 898. These films comprising a hydrocarbon
. · · · · · · · · · · · · · · · · · · ·	resin may be oriented in uniaxial or biaxial directions to the same or different degrees.
	the consequence of the problem is a best state of the contract
5	In a preferred embodiment this invention also relates to a method to produce a film
	characterized by good haze values comprising:
	i), selecting a first polymer having a CDBI of 50 % or more comprising
	homopolyethylene or a copolymer of ethylene and up to 50 weight % of a C ₃ to
	C_{20} olefin, where $s o s_{72}$ with the robbs burnes of the confidence s_{72}
10	ii) selecting a second polymer comprising homopolypropylene or a copolymer
	of propylene and up to 50 weight % of ethylene or a C ₄ to C ₂₀ olefin,
	iii) selecting a third polymer comprising one or more polymers produced in a
	high pressure process using a free radical initiator, and
	iv) combing the first, second and third polymers and forming them into a film.
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	In a preferred embodiment this invention also relates to a method of packaging an
	article comprising:
	i) selecting a first polymer having a CDBI of 50 % or more comprising
	homopolyethylene or a copolymer of ethylene and up to 50 weight % of a C ₃ to
20	Coolefin), is the above where a bias of the control in the first and the control in
	ii) selecting a second polymer comprising homopolypropylene or a copolymer
	of propylene and up to 50 weight % of ethylene or a C_4 to C_{20} olefin,
	iii) selecting optional polymers for core layers,
. 4	polymor
25	forms all or part of a film surface layer and the second polymer forms all or part
	of a film surface layer and, if present, the optional polymers for core layers are
	formed into film layers in between the first surface layer and the second surface
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	enclosing an article in the film; and bit follows are constructed by
30	vi) heat sealing the enclosed article such that at least one seal is formed by heat
	sealing the first surface layer to the second surface layer.
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	In a preferred embodiment the films formed from the blends described herein when
	formed into a film 50 µm thick film have an average secant modulus greater than 350
	MPa and a dart drop impact strength greater than 5 g/micron.
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Additionally the film produced herein can be laminated with another film such as polyethylene, polypropylene, polyester, polyamides and the like which may or may not be oriented. These combinations are particularly suitable for high quality packaging performance such as modified atmosphere packaging or controlled atmosphere; packaging: An Archive to the will be a fine to the archive to the

The films produced herein may be used for typical packaging applications, form fill and seal applications, cling films, stretch films, frozen film, heavy duty packaging film, can liners and other similar applications.

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ECD 109 is an ethylene hexene copolymer produced in the gas phase having about 4.1 weight % hexene, a melt index of about 0.8 g/10min, a CDBI of about 59, an Mw/Mn of about 2.3 and a density of about 0.928 g/cm³, sold under the 15 tradename EXCEEDTM by Exxon Chemical Company in Baytown, Texas.

ECD 103 is an ethylene hexene copolymer produced in the gas phase having (approximately 7.6 weight % hexene, a melt index of about Fg/10min, a Mw/Mn of about 2.3, a CDBI of about 67% and a density of about 0.917 g/cm³, sold under 20 the tradename EXCEED™ by Exxon Chemical Company in Baytown, Texas. on the card of the SC mergla 26 of the card of the card-

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ECD 202 is an ethylene hexene copolymer produced in the gas phase having approximately, 7.6 weight % hexene, a melt index of about 2.4 g/10min, a Mw/Mn of about 2.3, a CDBI of about 67% and a density of about 0.917 g/cm³, sold under the tradename EXCEED™ by Exxon Chemical Company in Baytown, Texas. the arriver of the property and the first of the property of t

PP-1 is a homopolymer of propylene having a Melt Index of about 2.9 g/10 min and a broad molecular weight distribution (Mw/Mn) sold under the trade name of 30 ESCORENE PP 4352F1 by Exxon Chemical Company

and without between which were a server will be given it

LD-2 is a low density polyethylene having a density of about 0.922 g/cm3 and a melt index of about 0.75 g/ 10 min commercially available under the trade name Geo and ESCORENE LD 150/BW from Exxon Chemical Belgium. The confidence of 大大大学 (1916年) 1916年 (1917年) 1918年 (1917年) 1918年 (1917年) 1918年 (1917年)

LL-3 is ESCORENE LLN 1201 XV a ethylene butene copolymer having a Melt Index of about 0.7 g/10 min, a density of 0.925 g/cm3 and produced in a gas phase

using a Ziegler Natta catalyst and is commercially available from Exxon Chemical Belgium.

TESTING METHODS:

Composition Distribution Breadth Index (CDBI) is measured by the procedure described in PCT publication WO 93/03093, published February 18, 1993.

Fractions having a molecular weight (Mw) less than 15,000 were ignored.

Melt Index (MI) was measured according to ASTM D 1238. (190 °C, 2.16 kg)

Density was measured according to ASTM D1505, where the sample was

Paramina de la

- prepared according to ASTM D 1928/.

 Mw and Mn were measured by GPC (Gel Permeation Chromatography) on a Waters
 150 gel permeation chromatograph equipped with a differential refractive index (DRI)
 detector and a Chromatix KMX-6 on line light scattering photometer. The system was
 used at 135 °C with 1,2,4-trichlorobenzene as the mobile phase. Shodex (Showa
- Denko America, Inc) polystyrene gel columns 802, 803, 804 and 805 were used. This technique is discussed in "Liquid Chromatography of Polymers and Related Materials III", J. Cazes editor, Marcel Dekker. 1981, p. 207, which is incorporated herein by reference. No corrections for column spreading were employed; however, data on generally accepted standards, e.g. National Bureau of Standards Polyethylene 1484 and
- anionically produced hydrogenated polyisoprenes (an alternating ethylene-propylene copolymer) demonstrated that such corrections on Mw/Mn (= MWD) were less than 0.05 units. Mw/Mn was calculated from elution times. The numerical analyses were performed using the commercially available Beckman/CIS customized LALLS software in conjunction with the standard Gel Permeation package. Calculations
- 25 involved in the characterization of polymers by ¹³CNMR follow the work of F. A. Bovey in "Polymer Conformation and Configuration" Academic Press, New York, 1969.
 - Dyna Impact Strength properties (Max Force, Damaging Energy, Total Energy,
- Damaging Travel, Total Travel) were measured according to DIN 53373.

 Tensile properties (Tensile at yield, elongation at yield, tensile at break, elongation at break, and Secant Modulus) were measured according to ASTM D 882. Average secant modulus is the mathematical average of the MD Secant Modulus and the TD Secant Modulus.
- 35 Elmendorf Tear Strength (N/μm) was measured according to ASTM 1922.

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Gloss was measured according to ASTM D2457/60° control of the participants.

Dart Dr p was measured acording to ASTM D 1709.

Heat seal testing procedure: Seal were made on a Topwave sealing machine. The film was folded between TEFLONTM film and inserted between the sealing bars. At various the sealing bars were closed with a pressure of 0.5 MPa for 0.5 seconds. The film was removed from the Top wave machine and conditioned for a minimum of 12 hours at 23 °C ± 3 °C and 50% humidity ± 5% humidity.

Seal Strength was tested according to the following procedure. After conditioning for a minimum of 12 hours at 23 °C \pm 3 °C and 50% humidity \pm 5% humidity, the seal

10 strength of 15mm wide sample was measured in a Zwick tensile instrument under the following conditions: speed-100 mm/min, load cell-200N, and clamp distance-50 mm.

The film was placed between the clamps and the clamps were moved apart at a speed of 100mm/min. During the rest the force (N) was recorded as a function of elongation (%). Four test specimens were measured and the average seal strength curve was recorded. The seal strength was the force at which the test specimen failed.

Example 16 to the control of the agency because the bond of the control of

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Eleven monolayer films 50 \(\mu\) thick of various blends were blown on an Alpine extruder under the conditions in Table 1, Table 3 and Table 6. The individual polymers were fed into the same extruder hopper at the same time. The blend components and test data on the resulting films are reported in Tables 2, 4, 5 and 7.

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Table 1

		Table			
하셔트 끝들은	ECD109 95w%	ECD109 90w%	ECD109 80w%	ECD109 80w%	ECD109 75w%
of the state	PP-1 5w%	PP-1 10w%	PP-1 20w%	PP-1 15w%	PP-1 20w%
	1000		. }	LD-2 5w%	LD-2 5w%
Barrell temp					,
settings (°C)	i			Í	:
zone 1	190	190	190	190	190
zone 2	200	200	200	199	200
zone 3	220;	220	220	220	220
zone 4	220	220	220	220	220
zone 6	230	230	225	235	230
zone 7	230	230	230	224	230
zone 8	230	230	222	235	230
zone 9"	240	240	239	241,	240
zone 10	240	240	- 240	240	240
zone 11	24ò	240	240	240	240
zone 12	240	240	240	240-0 - 0	240
diegap (mm)	1.5	1.5	1.5	1.5	1.5 for 1:5.
cooling air temp	16	,17		16	16 л
(°C)	'	d - contact and insurance	**************************************		- 1
Die Diam (mm)		200	200	200	200
Melt temp. (°C)	1.5 m	184	tigi		10 Apr 17
Ti	243	. 240		233	233
12	242	237	1	234	232
T3	245-	239	and the second of	237	235
T4	242	237		234	232
T5	245	242	The management of the second o	232	233
Tmelt	237	243		235 345	237
Melt pressure	1 761	1777	•	1,05 (2.00)	T.
(bar)		1111-001	Andrew Communication Control Communication C		
1	545	326	The second secon	464	,
P2	518	491	7 ·	444	· . · · .
P4	501	473 \$		432	
P5 · · · · · · · · · · · · · · · · · · ·	516	§ ₹ 500 ₹	& ·	480	
P6 .	388	545	Professional real of the parties and the second sec	524	
screw speed				339 1.	**
<u> </u>	59 rpm	60 rpm		62 rpm	· 62 rpm
Output (kg/hr)	140	140		140	140
Lay-flat (mm)	900	900		900	900
Frost line (mm)	700	700		700	700
take off(m/min)	28	28		28	28
Blow up ratio	2.9	2.9		2.9	2.9

w% = weight percent based upon the weight of the polymers.

Table 2:

20 11 11 19 19 19 19 19 19 19 19 19 19 19	ECD109	ECD109	ECD109	ECD109	ECD109
3.8 × × × × × × × × × × × × × × × × × × ×	95ŵ%	90w%	80w%	80w%	75w%
	PP-1	PP-1	PP-1	PP-1-	PP-1
	5w%	10w%	20w%	15 w %	,20w%
200		. ••		LD-	:LD-
	5	11.7%	_ '	2 5w% -	2:5w%
Haze (%)	19	``18	22	9.5	9.7
Gloss (%)	7.8	7.1	5,4	10.7	9.9
Max Force N/μ	1.3	955,1.3	1.3.2	1.2	1.2
Damaging Energy mJ/μ	20	96×17	1215	. 13	11
Total Energy mJ/µ	22	20	14	16	12
Damaging Travel (mm)	24	20	16	18	15
Total Travel (mm)	25	01128	18,	23 (.r>-17
Tensile @ Yield (Mpa) MD	14.1	16.2	18.7	16.1	
Elong @ Yield (%) MD	16,1	13.1	-12	16.8	14.1
Tensile @ Break (MPa) MD	61	64	62	61	60
Elong @ Break (%) MD	682	670	649	678	674
Energy MD (mJ/mm3)	168	₩31 7 7	176	. 180	1190
Tensile @ Yield (Mpa) TD	14.6	15.7	17.8	16.4	18.1
Elong @ Yield (%) TD	12;5	11.2	10.0	10.0	9.9
Tensile @ Break (MPa) TD	56	35c 57	50.	. 54	51
Elong @ Break (%) TD	724	č⊭1735	6982	719	706
Energy TD (mJ/mm3)	169	177	160	167	104
Secant Modulus (MPa) MD	335	444	548	. 442	539
Secant Modulus (MPa) TD	369	(QL440 ·	540 _c	478	557
Average Secant Modulus (MPa)	352	ET#442	544	460	⊕ 548
Elmendorf tear (g/µ) MD,	8.9	^{ਹਰ:} 7.8 ਂ	5.6	7.1	4.3
Elmendorf Tear (g/µ) TD	15	75 13 ·	10	16	15

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TD = tra	insve	rse direc	tion.	 	my 60
MD= m	achin	a directi	On.	 	CHI
MID= III	acmin	e difeen	OII. V	 	909
0.1		V/1		 	on:
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Table 3

		Table 3			
	ECD103			ECD103 75w%	
11 12	100 w%			PP-1 20w%	
		v 40	LD-2 5w%	LD-2 5w%	
Barrell temp		Sut in 1			
settings' (°C)	;				
zone i	170	190	190	191	
zone 2	175	200 ;	200	201	
zone 3-	175	220	220	219	
zone 4	175	220	220	- 219	
zone 6	185	230	225	224	
zone 7	185	230	230	230	
zone 8	185	230	222	228	
zone 9	190	240	239 -	240	
zone 10	190	240	240	240	
zone II	200	240	240	now 240 127	:
zone 12	200	240	240	240	Ett.
diegap (mm)	- :1:5: "	1.5	1.5	4.5	
cooling air temp	1			the state of the s	
(°C)	17,	16	16	57	
	200	200	200	200	٠. ·
Die Diam (mm)	200	200	1. 1. 200. L		 ! .
Melt temp. (°C)		243		230	 -
T1 37777	214 227	243	231 235	234	
T2	227	248		236	713
T3 ``	233 226	248	··· 238 ··· 纪 234	233	٠,٠
T4	*** *** *** ***		231	230	
T5	211	242	231	230	
Tmelt	206	>243	(in: 230		
Melt pressure	57	in ({-		
(bar)	446	\$200	276	245	
P1	440	\$0234	276	245	
P2	437.	55278	301]	
P3 ()	456	316	370	305	
P4	517	400	397	378	
P5	600.	465	446	425	
P6	414	326	304	288	
screw speed	52 rpm	68 rpm	67 rpm	68 rpm	
Output (kg/hr)	133	138	139	EN 2 141 .	
Lay-flat (mm)	930	900	:	900	
Frost line (mm): ,	2 0,600 F	17. 700 St. C.	700 💀 🖔		
take off(m/min)	26	28,	. 28	28	
blow up ratio	3.0	2.9	2.9	2.9	

w% = weight percent based upon the weight of the polymers.

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Table 4

and the party and the first and the party				·
	ECD103	ECD103	ECD103	ECD103
	100%	95w%	80w%	75w%
	a 4 (64 A P 1754)	LD 150	PP-1	PP-1
	·	5 w%	15w%	20w%
50,		3.3	LD-2	LD-2
e%		C.T	5w%	5w%
Haze (%)	25	5.4	3.7	4.7
Gloss (%)	5.8	ა 12.9	13.0	12.0
Max Force N/μ	>1.7	⊕ >1.7	1.4	1.4
Damaging Energy mJ/µ	>47	ː>47	24	22
Total Energy mJ/µ	>47	, .>47	26	25
Damaging Travel (mm)	>44	54°>44	26`	· 24 ///
Total Travel (mm)	>44	>44	28	27
Tensile @ Yield (Mpa) MD	10.0	10.2	_ 15.2	15.9
Elong @ Yield (%) MD	13.9	13.6	16	15.4
Tensile @ Break (MPa) MD	69	62	65,	64
Elong @ Break (%) MD	613	642	637	653
Energy MD (mJ/mm3)	143	148	173	178
Tensile @ Yield (Mpa) TD	9.7	10.5	13.6	14.3
Elong @ Yield (%) TD	20.0	31:16.6	12.0	12.0
Tensile @ Break (MPa) TD	. 60	Sect 59	56	56
Elong @ Break (%) TD	661	671	692	
Energy TD (mJ/mm3)	140	144	157	158
Secant Modulus (MPa) MD	. 183	heg202	395	; 443 :
Secant Modulus (MPa) TD	189	and221	361	394
Average Secant Modulus (MPa)	186	4. 212 i	378	419
Elmendorf tear (g/µ) MD	12	12 :	13	12
Elmendorf Tear (g/µ) TD	14	JE 17	16	16

Table 5 Heat Seal Strength (N/15mm) of 50 micron films

	Table of Izent Sent Att on Girls (1.1.29, Inn.) of Sent Inn.						
Heat Seal Temp	ECD 103 95 w%	ECD 103 75 w%	ECD 109 75 w%				
f	LD-2 5 w%	PP-1 20 w%	PP ₇ 1 20 w%				
1	·	LD-2 5w%	LD-2 5w%				
110 °C	6.0	0.7					
120°C	9.5	11.5	1.2				
130°C	10.0	12.3	13.1				
140°C	9.2	11.3	13.4				
150°C	11.1	11.0	13.1				
160°C	12.0	10.7	12.5				
180°C	11.7	11.0	12.6				

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	Table 6	et vent - tomos - e -				
WAY BUILD OF BUILD	LL-3'80w%	ECD202 80w%				
	LD-150 20w%	PP-1 15w%				
		LD-2 5w%	. ;			
Barrell temp settings (°C)					
zone 1	170	170				
zone 2	175	175				
zone 3	175	180				
zone 4	175	190				
zone 6.	185	190				
zone 7:1	185	190				
zone 8	190	195). j.			
zone 9	190	215	(:			
zone 10	190	215	+ 4) 1			
zone 1·1	200	225				
zone 12	200	225	- 44 A			
diegap (mm)	1.5	1.5				
cooling air temp (°C)		15				
Die Diam (mm)	1 5 5 5 5 1 200 Hz 11 1 1 1	0 7 200 0 1 Char	.•			
Melt temp. (°C)	Country Mines in	The first of the little	554			
$\frac{\Gamma_1}{\Gamma_2}$. The two common left gives	107	197				
	205	205				
T3 : Sandilities and only		219	•			
T4	204	204 4				
T5	196	196				
Tmelt	196	196				
Melt pressure (bar)						
P1	563	201				
P2	620	207				
P3	595	210				
P4	556	207				
P5	552	200				
P6	412	200				
screw speed	41 rpm	44 rpm				
Output (kg/hr)	100	95				
Lay-flat (mm)	785	785				
Frost line (mm)	500	500				
take off(m/min)	50	62				
blow up ratio	2.5	2.5				

w% = weight percent based upon the weight of the polymers.

Гa	hl	e.	7

25 75 S023 32	: \	3.80w%	ECD202 80w%
(3.1) 2.57		LD-150 20w%	PP-1 15w% ;
			LD-2 5w%
Haze (%)	1	2.7	3.8
Gloss (%)	۲.	13.6	12.2 :
Tensne @ Break (MPa) MD	1	68.1	98.1 a 📑
Elong @ Break (%) MD	4	474	552
Tensile @ Break (MPa) TD	873	35.4	49.4 🕮 🔩
Elong @ Break (%) TD	:31	815	695° ±00°
Secant Modulus (MPa) MD	3,53	407	337 \$.50
Secant Modulus (MPa) TD		502	316 936
Dart Drop (g/μm)	4,500	3.1	8.1
Elmendorf tear (g/µ) MD	(t)	0,3	4.0
Elmendorf Tear (g/μ) TD	.4.7	13.7	15,7

All documents described herein are incorporated by reference herein, including any priority documents and/or testing procedures. As is apparent from the foregoing general description and the specific embodiments, while forms of the invention have been illustrated and described, various modifications can be made without departing from the spirit and scope of the invention. Accordingly, it is not intended that the invention be limited thereby.

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What is claimed is: Managed to Alexander of John Alexander

14.1. A film comprising a blend comprising:

- (i) a homopolymer of ethylene having an Mw/Mn of 3 or less or a copolymer of ethylene and up to 50 weight% of a C₃ to C₂₀ olefin having a CDBI of 50 % or more,
 - (ii) a homopolymer of propylene or copolymer of propylene and up to 50 weight % of a comonomer, and an arrange of the strength of the little of the strength o
 - (iii) a polymer produced in a high pressure process using a free radical initiator.

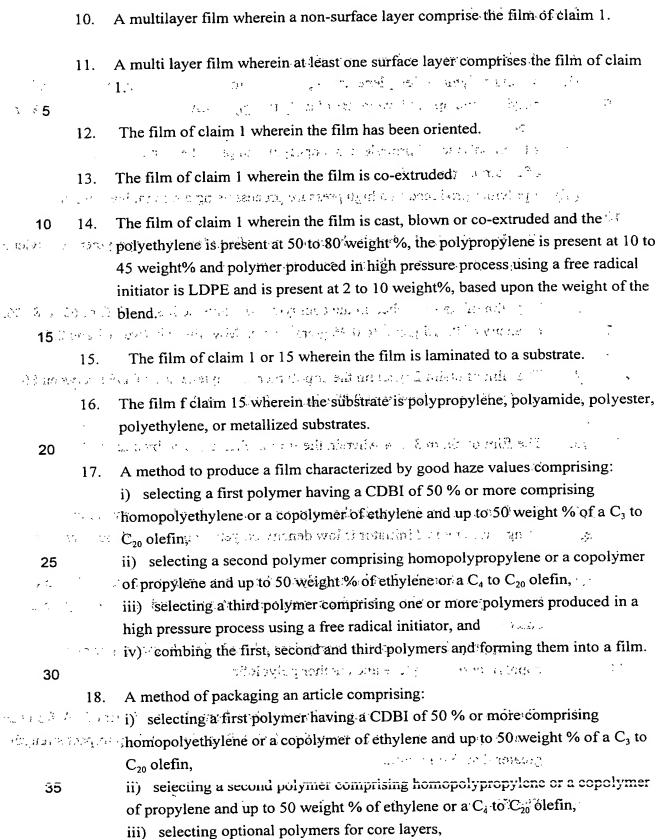
10 and combine with a complete consist of millions in the second of a million of the consist of

- The film of claim 1 wherein the copolymer of ethylene is a copolymer of ethylene and propylene, butene, hexene and/or octene: A fine of the film of ethylene and or octene.
 - 3. The film of claim 2 wherein the copolymer of ethylene has a CDBI of 65 to 85 %, a density of 0.915 g/cm³ to 0.96 g/cm³ and an Mw/Mn of between 1 and 2.5?
- - 20 5. The film of claim 3 or 4 wherein the film is characterized by a haze of 5 % or the start of classifier over the gradient measurement of the start of the st
 - 16. The film of claim lawherein the polymer produced in a high pressure process using a free radical initiator is low density polyethylene or ethylene vinyl acetate.
 - 25 Committee of the com
 - 7. The film of claimal wherein the polyethylene component comprises a blend of the homopolymer of ethylene or the copolymer of ethylene and another polyolefin, and/or and solve the polypropylene component comprises a blend of the homopolymer or copolymer of propylene and another polyolefin.
 - 8. The film of claim 1 characterized in that when formed into a 50 µm thick film has we an average secant modulus greater than 350 MPa and a dart drop impact strength greater than 5 g/micron.

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9. The film of claim 1 wherein the film is sealed.

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- iv) combing the first polymer and second polymer so that the first polymer forms all or part of a film surface layer and the second polymer forms all or part of a film surface layer and, if present, the optional polymers for core layers are formed into film layers in between the first surface layer and the second surface layer,
- v) enclosing an article in the film, and

vi) heat sealing the enclosed article such that at least one seal is formed by heat sealing the first surface layer to the second surface layer....

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INTERNATIONAL SEARCH REPORT

1205

Intr I onal Application No
PC1/US 98/03597

A. CLASSIF	FICATION OF SUBJECT MATTER C08L23/10 C08L23/08 C08L23	1/04 C08.15/18 B3	32B27/32	
	COOLEST TO COOLEST OF COOLEST			
According to	International Patent Classification (IPC) or to both national class	ification and IPC	g . * * * * * * * * * * * * * * * * * *	
B. FIELDS	SEARCHED CO. S. A. A. A. A. A. A. S. S. S. M. M. C.	m. To the configuration of	į. į	
Minimum do IPC 6	cumentation searched (classification system followed by classific COSL COSJ B32B	ation symbols)	erio. Testa	
Documentati	ion searched other than minimum documentation to the extent tha	at such documents are included in the field	s searched	
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Electronic da	ata base consulted during the International search (name of data	base and, where practical, search terms of	used)	
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.	
Р,Х	WO 97 28960 A (TREDEGAR IND INC 1997	2) 14 August	1-18	
	see page 1, line 27 see page 2, line 4 - line 27 see page 4, line 16 - line 27 see page 6, line 21 - page 7, l	ine 19		
X	US 5 424 362 A (HWANG YUH-CHIN June 1995 see column 7, line 39 - line 42		1-9,17	
χ	WO 92 14784 A (EXXON CHEMICAL F	PATENTS) 3	18	
Y	September 1992 see page 3, line 28 - line 29; example IX		1-18	
		-/		
X Furt	her documents are listed in the continuation of box C.	Patent family members are li	isted in annex.	
° Special ca	ategories of cited documents :	"T" later document published after the or priority date and not in conflic	t with the application but	
consid	ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international date	cited to understand the principle invention "X" document of particular relevance cannot be considered novel or c	; the claimed invention cannot be considered to	
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or				
other	means ent published prior to the international filing date but	ments, such combination being in the art.		
	actual completion of the international search	Date of mailing of the internation		
2	26 May 1998	08/06/1998	·	
Name and	mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk	Authorized officer		
	Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Clemente Garci	a, R	

INTERNATIONAL SEARCH REPORT

ory ,	Citation of document, with indication where appropriate, of the relevant passages	Relevant to claim No.:		
	WO 94 26816 A (EXXON CHEMICAL PATENTS INC.) 24 November 1994 cited in the application see page 19, line 11 Jine 29; claims	1-18		
	中では、	MARIO LA SEMINISTA		
		0.80 <u>8</u> 0202 €		

Form PCT/ISA/210 (continuation of second sheet) (July 1992)

INTERNATIONAL SEARCH REPORT

Information on patent family members

Int ional Application No PCI/US 98/03597

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